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TITLE: Anthelmintic 5-substituted aminobenzimidazoles

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FR 1580153 GB 1198941			FR GB	
GB 1198942			GB	
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GI For diagram(s), see printed CA Issue.

The title compds. (I) are anthelmintics, particularly useful in treating AB veterinary nematode infections (given orally) and also possess antifungal properties. To a solution of 10 g II (R1 = H, R2 = 4-thiazoly1) in 20 ml concentrated H2SO4 was added dropwise at 20-30° a mixture of 4 ml concentrated HNO3and 6 ml concentrated H2SO4, and the mixture kept 5 min to give II (R1 =

02N.

R2 = 4-thiazolyl) (III), m. 240-1° (HCONMe2) (DMF). A solution of 2.2 g furfural in 3 ml EtOH was added to 3 g 3,4-(H2N)2C6H3-NO2 (IV) suspended in 10 ml PhNO2, and the mixture stirred 10 min at room temperature then heated slowly to 210° 1 min (allowing the EtOH to dist.) to give II (R1 = O2N, R2 = 2=furyl), m. 224°. II (R1 = O2N, R2 = 2-pyrryl), m. 259-60° was prepared by refluxing a mixture of 43.2 g 2-formylpyrrole, 54 g IV, and 160 g Cu(OAc)2 in 1 1. MeOH 2 hr. Catalytic hydrogenation of 141 g III in 4 1. EtOH over 22 g 5% Pd-C at 24° and 45 psi 5.5 hr gave II (R1 =H2N, R2 = 4-thiazolyl) (V), m. 232-3° (EtOH-Hexane). Addition of 1 g ClCO2Me to a suspension of 2.16 g V in 7.5 ml C5H5N gave II (R1 = MeO2CNH, R2 = 4-thiazolyl) (Va), m. 237-9° (MeOH containing C), also obtained in the absence of C5H5N using Me2CO as solvent at room

temperature

Similarly were prepared the following II (R2 = 4-thiazolyl) (R1 and m.p. given): EtO2CNH, 203-5° (MeCN-Et20) (methanolate m. 94-105°); PrO2CNH, 214-15°; BuO2CNH, 211-12°; C5H1102CNH, 178-9°; C6H1302CNH, 150-2°; n-C8H1702CNH, 66-7°; PhO2CNH, 115-16°; 4-FC6H4O2CNH, 275-80°; 2-FC6H4O2CNH, 135-40°; iso-BuO2CNH, 231-2°; Me2CHO2CNH,

212-14°; CH2:CHCH2O2CNH, 210-12°; HC.tplbond.CCH2O2CNH,

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200-2°; EtSC(0)NH, 215°; and cyclopropyloxycarbonylamino,
     190-5° (hydrate), 207-8° (anhydrous). By standard acylation
     procedures were prepared the following I (R1 = H, X = O), (R2 and m.p.
     given): Me, 280°; PhCH2, 210-11°; H, 247-8°; Et,
     255-6°; Ph (VI), 118-20°; 3-pyridyl, 284-5°; 2-FC6H4,
     132-3°; 1-adamantyl, 246-7°; 2-naphthyl, 154-6°;
     cyclopropyl, 245°; Me2CH, 203-5°; 3-thienyl, 276-8°;
     3-FC6H4, 232-3°; 4-FC6H4, 151-2°; 2-MeOC6H4, 113-14°;
     3-MeOC6H4, 105-9°; 2-PhO, 95-100°; 2-C1C6H4, 146-7°;
     3-IC6H4, 127-9°; 3-F3CC6H4, 201-3°; 3-O2NC6H4,
     163-4°; 2,5-F2C6H3, 113-14°; 2-pyridyl, 240-1°;
     4-pyridyl, 150-3°; Me3C, 241-2°; 2-furyl, 139-40°;
     4-thiazolvl, 387-8°; 2-thienvl, 288° (decomposition); MeOCH2,
     238-9°; C12CH, 220°; and Me2C:CH, 270-2°. Refluxing
     a mixture of 5 g I (R1 = H, R2 = EtS, X = O), (VIa) 0.5 g Bu2SnO, and 50 ml
     F3CCH2OH 20 hr gave I (R1 = H,R2 = F3CCH2O), m. 231-2°
     (AcOEt-hexane). I (R1 = H, R2 = HC.tplbond.CCH2O, X = O), m.
     202-2°, was similarly prepared Addition of 3.62 g PhOC(S)Cl dropwise to
     4.32 \text{ q V} in 25 \text{ ml C5H5N} and stirring 1.5 \text{ hr} gave I (R1 = H, R2 = PhO, X =
     S), m. 155-7° (MeOH), which on heating in C5H5N (7 vols.) 1 hr at
     100° gave 5-isothiocvanato-2-(4-thiazolv1)benzimidazole (VII), m.
     243-6^{\circ}. I (R1 = H, R2 = MeO, X = S), m. 224^{\circ} (MeOH), was
     prepared by refluxing a mixture of 2.5 g VII, 25 mg NaOMe, and 300 ml MeOH20
     hr; I (R1 = H, R2 = EtO, X = S), prepared similarly, m. 218°. A slow
     stream of MeSH was passed into 4 q VII in 25 ml Me2NCHO 15 min, and the
     mixture kept 20 hr to give I (R1 = H, R2 = MeS, X = S), m, 202-5°. I
     (R1 = H, R2 = Ph, X = S), m. 140-3° (MeOH), was obtained by
     refluxing a mixture of 1 g VI, 2 g P2S5, and 20 ml C5H5N 25 min. A
     suspension of V (amount unspecified) in 25 ml C5H5N was treated dropwise
     with 1.2 g MeNCO and the mixture stirred 2.5 hr to give I (R1 = H, R2 =
     MeNH, X = 0), m. 160° (MeOH). Dropwise addition of 2.5 g Me2NCOC1 to
     a suspension of 4.32 	ext{ g V} in 25 	ext{ ml C5H5N gave I (R1 = H, R2 = Me2N, X = O),}
     m. 260-2° (MeOH). Refluxing a mixture of 5 g VIa and 25 ml Et2NH 6
     hr gave I (R1 = H, R2 = Et2N, X = O), m. 234-5^{\circ}. The following I
     (R1 = H) were similarly prepared (R2, X, and m.p.given): pyrrolidino, O,
     296-8°; piperidino, O, -; MeNH, S, 235-7°; PhNH, S,
     244-6°; Me2N, S, 156-9°; Et2N, S, 130-5°; piperidino,
     S, 225-6°, and pyrrolidino, S, 257-8°. A mixture of 8.5 q I
     (R1 = H, R2 = Me2CHO, X = O) (VIII), 100 ml DMF and 2.3 q 52% NaOH
     emulsion in mineral oil was stirred at room temperature 20 min., heated 10 min
     at 50°, cooled to room temperature, treated with 7.1 g MeI in 10 ml DMF,
     and the mixture heated 20min at 80° to give I (R1 = Me, R2 = Me2CHO,
     X = O), purified by crystallization from AcOEt. Addition of a mixture of 1.3
ml concentrated
     HNO3 (d. 1.41) and 2.8 ml concentrated H2SO4 to 3.8 g 1-methoxy-2-(4-
     thiazolyl)benzimidazole in 12.3 ml concentrated H2SO4 at 12° gave 1.5 g
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ncentrated HNO3 (d. 1.41) and 2.8 ml concentrated H2SO4 to 3.8 g 1-methoxy-2-(4-thiazoly1)benzimidazole in 12.3 ml concentrated H2SO4 at 12° gave 1.5 g 5-nitro-1-methoxy-2-(4-thiazoly1)benzimidazole (IX). m. 220-1° (MeOH). Hydrogenation of IX in MeOH and treatment with methanolic HCl gave 450 mg amorphous 5-amino analog-HCl, which with ClCO2CHMe2 in CSHSN gave I (RI = MeO, R2 = MeZCHO, X = 0), m. 123-5°. Treatment of 5.4 g Va in 100 ml PhMe and 30 ml DMF with 0.7 g NaH in 2 ml PhMe at 65°, stirring 1 hr, adding 2.5 g ACCl at 55°, and refluxing 0.5 hr gave I (RI = AC, R2 = MeO, X = 0). A mixture of 3.26 g VIII and 1 g BUNCO in 100 ml MeCN was refluxed 4 hr to give I (RI = BUNHCO, R2 =

Me2CHO, X=0). Treatment of VIII in C5H5N with ClCO2CHMe2 gave I (R1 = Me2CHO2C, R2 = Me2CHO, X=0), crystallized from aqueous MeOH. Similarly, V

and

MeSO2Cl gave II (R1 = MeSO2NH, R2 = 4-thiazolvl), m. 225-6° (MeOH). II (R1 = PhSO2NH, R2 = 4-thiazolyl) (X), m. 254-5°, was also prepared Methylation of 3.5 g X in 10 ml MeOH containing 0.54 g NaOMe with 0.625 ml MeI gave II (R1 PhSO2NMe, R2 = 4-thiazolyl), m. 142-3° (MeOH); II (R1 = MeSO2NMe, R2 = 4-thiazolyl), m. 192-3° (MeCN), was also prepared The following XI were analogously prepared (R1, R2, R3, and m.p. given): H, MeO2CNMe, 4-thiazolvl, 161-2°; H, MeO2CNH, 2-furvl, 162-3°; MeO, MeO2CNH, 2-furyl, 164°; H, EtO2CNH, 2-furyl, 171-2°; H, PhO2CNH, 2-furyl, 150-5°; H, EtO2CNH, 2-pyrryl, 200-2°; H, MeO2CNH, 2-thienyl, 185-8°; H, MeO2CNH, 1,2,5-thiadiazol-3-yl, 150-5°; H, MeO2CNH, 1-pyrazolyl, 207-10°; H, MeO2CNH, 2-methyl-4-thiazolyl, 135°; H, MeO2CNH, 1,2,3,-thiadiazol-4-yl, 218-20°; H, MeO2CNH, 1,3,4-thiadiazol-2-vl, 258°; H, Me2CHO2CNH, 2-oxazolyl, 206°; H, Me2CHO2CNH, 2-thiazolyl, 234°; H, MeO2CNH, 2-imidazolyl, 205-7°; H, 4-FC6H4CONH, 2-furyl, 264°; H, 2-furoylamino, 2-furyl, 248°; H, 4-FC6H4CONH, 1-pyrazolyl, 230°; H, BzNH, 2-thiazolyl, 135-40°.

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 27217-07-8 CAPLUS

CN Urea, N,N-dimethyl-N'-[2-(4-thiazolyl)-1H-benzimidazol-5-yl]- (9CI) (CA INDEX NAME)

Me2N-C-NH

RN 27217-08-9 CAPLUS

CN Urea, 1,1-diethyl-3-[2-(4-thiazolyl)-5-benzimidazolyl]- (8CI) (CA INDEX NAME)

Et2N-C-NH

RN 27217-09-0 CAPLUS

CN 1-Pyrrolidinecarboxamide, N-[2-(4-thiazoly1)-5-benzimidazoly1]- (8CI) (CA INDEX NAME)

- RN 28767-12-6 CAPLUS
- CN Urea, 1-methyl-3-[2-(4-thiazolyl)-5-benzimidazolyl]- (8CI) (CA INDEX NAME)